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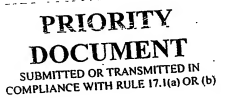
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The attached documents are exact copies of the European patent application described on the following page, as originally filed.

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Patentanmeldung Nr. Patent application No. Demande de brevet n°

03020119.8



Der Präsident des Europäischen Patentamts; Im Auftrag

For the President of the European Patent Office

Le Président de l'Office européen des brevets p.o.

R C van Dijk

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Process for the preparation of (meth)acrylate di-ammonium salts and their use as monomers for the synthesis of polymers

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22.35.EP

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Process for the preparation of (meth)acrylate di-ammonium salts and their use as monomers for the synthesis of polymers

The invention relates to the preparation of (meth)acrylate di-ammonium salts of high purity and their use as monomers for the synthesis of polymers useful as cationic flocculants in water treatment.

(Meth)acrylate ammonium salts and their use as monomers for the synthesis of polymers useful as flocculants have been described in WO 01/55088, WO 01/55225 and US2002/0035198. In these publications the (meth)acrylate ammonium salts are synthesised by the reaction of the corresponding di-amine (meth)acrylate with less than 2 equivalents of an alkyl or a benzylhalide in chloroform and in water. These processes yield aqueous solutions of a mixture of a (meth)acrylate di-ammonium salt and a mono-amino (meth)acrylate mono-ammonium salt. These processes do not permit to isolate (meth)acrylate di-ammonium salts with a high purity. It is also almost impossible to purify the obtained mixtures. When used as monomers in the synthesis of polymers, these polymers will inevitably contain a significant amount of units derived from the mono-ammonium salts. In order to be useful as starting material for the production of high molecular weight polymers and copolymers for flocculation, the di-ammonium salts need to be very pure.

The present invention now provides a process for the manufacture of (meth)acrylate di-ammonium salts that overcomes the above-mentioned problems.

The present invention therefore relates to a process for the manufacture of a (meth)acrylate di-ammonium salt of formula (I)

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wherein \mathbb{R}^1 represents hydrogen or methyl, each \mathbb{R}^2 , independently, represents an alkyl comprising from 1 to 4 carbon atoms, each \mathbb{R}^3 , independently, represents an alkyl or an aralkyl and each X^* , independently, represents an anion, comprising (a) the reaction of the di-amino-(meth)acrylate of formula (II)

with at least 2 equivalents of at least one alkyl or aralkyl compound of formula \mathbb{R}^3X in an organic-solvent containing at most 5000 ppm of water and wherein the diammonium salt of formula (I) has a solubility at 25 °C of less than 1 g/100 g of solvent and wherein the solubility of the corresponding amino-(meth)acrylate ammonium salt of formula (V)

$$H_{2}C \xrightarrow{R^{1}} C \xrightarrow{Q} C \xrightarrow{R^{2}} C \xrightarrow{R^{2$$

has a solubility at 25 °C of at least 20 g/100 g of solvent; and (b) the separation of the di-ammonium salt of formula (I) from the reaction mixture

without dissolving it in water.

The term "alkyl", as used herein, is defined as including saturated, monovalent hydrocarbon radicals having straight, branched or cyclic moieties or combinations thereof.

The term "aralkyl", as used herein, is defined as a radical of formula $-(CH_2)_{11}$ -aryl 15 wherein n is an integer from 1 to 4 and aryl means any aromatic hydrocarbon having 6 to 24 carbon ring atoms that may be monocyclic or annealed. In this specification the term "(meth)acrylate" means "acrylate" as well as "methacrylate".

In the process according to the present invention, the organic solvent used in step (a) preferably contains at most 1000 ppm of water. The organic solvent used is 20 preferably a solvent wherein the solubility of the di-ammonium salt of formula (1) has a solubility at 25°C of less than 0.5 g/100 g of solvent.

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In the process according to the present invention, the solvent used in step (a) is usually an aprotic dipolar solvent; preferably acetone, methylethylketone, ethylacetate, nitromethane, acetonitrile or mixtures thereof. Particularly preferred is acetonitrile.

In the process according to the invention, step (a) is preferably carried out at a temperature ranging from 40 to 100 °C, most preferably from 70 to 90 °C. The process is preferably conducted at autogenic pressure in a closed reactor.

In the process according to the invention, step (a) is advantageously conducted with a molar ratio of the alkyl or aralkyl compound of formula R³X to the di-amino-(meth)acrylate of formula (II) higher than 2, most preferably of at least 2.1. The molar ratio preferably does not exceed 4.5, most preferably it does not exceed 3.

The duration of step (a) is generally from 1 to 100 hours, preferably from 10 to 30 hours.

The separation of the (meth)acrylate di-ammonium salt of formula (I) in step (b) of the process according to the invention may be carried out by any means suitable therefore. It is advantageously done by the filtration of the reaction mixture.

In an embodiment of the process according to the invention, the excess of alkyl or aralkyl derivative of formula R³X used in step (a) is separated from the reaction mixture, for example by stripping, before effectuating the separation in step (b).

The process according to the invention can be done as batch or continuously. In the latter case, the (meth)acrylate di-ammonium salt of formula (I) formed during step (a) can be separated from the reaction mixture continuously, for example by filtration, decantation or any other mean suitable therefore, and the reaction mixture can then be recycled and used as solvent in a subsequent reaction step (a).

According to another preferred embodiment of the process, the reaction mixture obtained after step (b) is recycled. In this embodiment, it is particularly preferred that the reaction mixture obtained after step (a) is filtered in step (b) in order to separate the (meth)acrylate di-ammonium salt of formula (i) already formed and to recycle the filtrate in a subsequent step (a) in order to continue the reaction. This recycling operation can be repeated several times. The recycling of the filtrate permits to increase even more the yield and the purity of the (meth)acrylate di-ammonium salt of formula (i).

In a variant of the process according to the invention, both embodiments are combined.

The di-amino (meth)acrylate of formula (II) used in step (a) of the process according to the invention can be obtained by any process suitable therefore. It is preferably obtained by the transesterification of a 1,3-di-amino-2-propanol of formula (III)

wherein each \mathbb{R}^2 , independently, represents an alkyl comprising from 1 to 4 carbon atoms, with a (meth)acrylate of formula (IV)

wherein R⁴ represents an alkyl comprising from 1 to 4 carbon atoms, in the presence of a lithium-based catalyst. It was found that the transesterification conducted with the aid of a lithium-based catalyst permitted to obtain high purity di-amino-(meth)acrylates with high yields. The synthesis of di-amino-(meth)acrylate of formula (II) has already been described in US3586711, in FR1568382, in FR1529000, in Zh.Org.Khim. 1969, 5(11), p.1947-1952, and in US 2002/183543. However, it was found that the process described in these documents gave relatively moderate yields and insufficient purity. Moreover, the reaction temperatures necessary for conducting these processes were high so that often uncontrolled side-reactions and polymerisation occurred. It was surprisingly found that using a lithium-based catalyst could overcome these problems.

The present invention therefore also relates to a process for the manufacture of a di-amino-(meth)acrylate of formula (II) by the transesterification of a 1.3-di-amino-2-propanol of formula (III)

Empf.nr.:090 P.014

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wherein each R², independently, represents an alkyl comprising from 1 to 4 carbon atoms, with a (meth)acrylate of formula (IV)

$$H_2C = C$$
 $O = R^4$
 $O = R^4$

wherein R⁴ represents an alkyl comprising from 1 to 4 carbon atoms, in the presence of a lithium-based catalyst.

In this process for the manufacture of a di-amino-(meth)acrylate of formula (II), the lithium based catalyst is preferably chosen from lithium oxide (Li₂O), lithium hydroxide (LiOH), lithium carbonate (Li₂CO₃), lithiumalkoxides such as methoxylithium (LiOCH₃), tertiobutoxylithium (LiOtBu), the lithium salt of a 1,3-dialkyl amino-2-propanol corresponding to formula (III), lithium citrate, lithium chloride (LiCl), Li-stearate (LiC₁₈H₃₅O₂), LiClO₄, Li₂SO₄, LiOAc, LiOOCPh and/or lithium bromide (LiBr) and their mixtures. Especially preferred are lithiumoxide, lithiumhydroxide and lithiumalkoxides, especially lithium methoxide, and their mixtures.

In this transesterification process, the lithium-based catalyst is generally used in an amount of 1 to 20, preferably in an amount of 4 to 10, equivalents of Li per mole of 1,3-di-amino-2-propanol of formula [III].

In this transesterification process, the temperature is preferably not exceeding 120 °C, more preferably the temperature is lower than 110 °C. The transesterification is advantageously carried out at a temperature of at least 80 °C.

The transesterification process is generally conducted at a pressure adapted to the desired reaction temperature.

The transesterification process is preferably carried out in the presence of from 500 to 3000 ppm (relative to the total weight of reaction mixture) of at least one stabiliser, preferably selected from radical inhibitors such as quinones, hydroquinones, phenothiazine, tris(nonylphenyl)phosphite. Preferred stabilisers are methyl ether of hydroquinone, phenothiazine, tris(nonylphenyl)phosphite, N.N-diphenyl-1.4-phenylenediamine, ethylenediaminetetraacetic acid and their mixtures.

The transesterification process is generally carried out with a (meth)acrylate of formula (IV) to 1,3-di-amino-2-propanol of formula (III) molar ratio of from 1 to 10, preferably from 3 to 6.

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The transesterification process is preferably conducted by introducing progressively the di-amine of formula (III) to the reaction mixture containing the catalyst and the (meth)acrylate of formula (IV) so that the formation of side-products is even more reduced.

(Meth)acrylates of formula (IV) wherein R4 is methyl or ethyl, especially methyl, are preferred.

The di-amine (meth)acrylates of formula (II) obtained after the transesterification reaction are preferably isolated from the reaction mixture by distillation, more preferably under vacuum, and most preferably after removal of the lithium-based catalyst, for example by filtration and/or adsorption on silica.

The di-amino-(meth)acrylates of formula (II) are preferably stabilised by the addition thereto of less than 500 ppm of one or more stabilisers as described here above.

In the process according to the invention, compounds of formula (I). (II) and (IV) wherein R¹ is methyl are especially preferred.

In the process according to the invention, compounds of formula (i), (ii) and (iii) wherein \mathbb{R}^2 is methyl are especially preferred.

In the process according to the invention. (meth) acrylate di-ammonium salts of formula (I) wherein each \mathbb{R}^3 , independently, is an alkyl comprising from 1 to 4 carbon atoms or benzyl, are preferred. Most preferred are compounds wherein each \mathbb{R}^3 , independently, is methyl or benzyl; especially methyl. (Meth) acrylate di-ammonium salts of formula (I) wherein both \mathbb{R}^3 are the same are preferred.

In the process according to the invention, (meth) acrylate di-ammonium salts of formula (I) wherein each X, independently, is an anion selected from halides, especially chloride and bromide, and methylsulfonates are preferred. Especially preferred is chloride.

The process according to the invention permits to obtain a high yield of the desired (meth)acrylate di-ammonium salts of formula (f). Yields of at least 90, even 99. % can be obtained, resulting in 99+ pure [meth)acrylate di-ammonium salts of formula (f). Almost no side products are formed: In the final product the amount of impurities such as alkylhalogenides, alcohol and acids that could disturb further polymerisation of the product, are very low. The quantity of amino-(meth)acrylate ammonium salt of formula (V) (relative to the amount of di-ammonium salt of formula (I)) in the final product is very low, usually less than 10 mole %, preferably less than 5 mole %.

Moreover, the process permits to obtain the products in pure solid form. In this form the products show high stability and can be stored for long periods without

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decomposition. Once the solid product is isolated aqueous solutions and formulations with other monomers can be made as required for the envisaged application.

The present invention therefore also relates to (meth)acrylate di-ammonium salts of formula (I) comprising less than 10 mole % (relative to the amount of (meth)acrylate di-ammonium salt of formula (II) of the corresponding amino-(meth)acrylate mono-salt of formula (V). The quantity of amino-(meth)acrylate mono-salt of formula (V) is preferably less than 5 mole %, most preferably less than 1 mole%.

When used for the manufacture of polymers it is preferred to use products as pure as possible so that the nature of the polymer obtained can be controlled, and also to avoid the presence of impurities that can disturb the polymerisation. It has been found that the (meth)acrylate di-ammonium salts according to the present invention satisfy these requirements. The present invention therefore also relates to the use of (meth)acrylate di-ammonium salts according to invention for the manufacture of polymers comprising less than 10 mole % preferably less than 5 mole %, more preferably less than 1 mole%) of units derived from an amino- (met)acrylate ammonium salt of formula (V) and to the polymers which can thereby be obtained.

The present invention therefore relates also to a polymer obtained from a monomer composition comprising, per 100 parts by moles.

(a) from 1 to 100 parts, preferably from 2 to 70 parts, by moles of at least one (meth)acrylate di-ammonium salt of formula (I) according to the present invention.

(b) from 0 to 99 parts, preferably from 30 to 98 parts, by moles of at least one acrylamide monomer of formula (VI)

$$H_2C = C - C - N = R^5$$
(VI)

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wherein R⁵ is hydrogen or methyl R⁶ and R⁷ are, independently, hydrogen, alkyl comprising from 1 to 6 carbon atoms, optionally substituted by one or more hydroxy or alkoxy groups:

and less than 10 mole % relative to the amount of (meth)acrylate di-ammonium salt of formula (I), of amino-(meth)acrylate ammonium salt of formula (V).

The acrylamide monomer of formula (VI) is preferably acrylamide.

The polymers according to the invention may further comprise:

[c) from 0 to 60 parts by moles of at least one water-soluble monomer which is potentially anionic by varying the pH, and which is preferably chosen from ethylenically unsaturated carboxylic acids and salts thereof and ethylenically

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unsaturated sulphonated monomers and salts thereof (such as acrylic acid, 2-acrylamido-2-methylpropanesulfonic acid and salts thereof);

(d) from 0 to 90 parts by moles of at least one cattonic water-soluble monomer of formula (VII)

wherein R⁸ is hydrogen or methyl, A is -O or NH-. B is -CH₂-CH₂-, -CH₂-CH₂-CH₂-or -CH₂-CHOH-CH₂-, R⁹ and R¹⁰ are, independently, an alkyl comprising from 1 to

16 carbon atoms, R¹¹ is hydrogen or an alkyl comprising from 1 to 16 carbon atoms, and Y is a monovalent anion (such as (meth)acrylamidopropyltrimethylammonium and (meth)acryloyl-oxyethyltrimethylammonium halides);

(e) from 0 to 10 parts by moles of at least one hydrophobic monomer, preferably chosen from alkyl(meth)acrylates and vinylaromatic monomers (such as ethylacrylate, butylacrylate, styrene), and/or

(f) from 0 to 30 parts by moles of at least one water-soluble monomer other than (a), (b). (c) and (d), preferably chosen from polyethoxylated (meth)acrylates, polyethoxylated (meth)acrylates containing hydrophobic units or aryl units and N-vinylpyrrolidone.

The polymers according to the invention preferably contain less than 5 mole %, more preferably less than 1 mole %, relative to the amount in moles of (meth)acrylate di-ammonium salt of formula (i). of amino-(meth)acrylate ammonium salt of formula (v).

Based on the (meth)acrylate di-ammonium salts according to the invention, polymers and copolymers of high molecular weight can be obtained, for example by solution polymerisation or polymerisation in dispersed media (emulsion and suspension). Preferred ways for preparation of such polymers are inverse emulsion and suspension polymerisation processes. By inverse macroemulsion polymerisation, high molecular weight cationic acrylamide-based co- and terpolymers with different charges can be obtained.

It was found that the polymerisation rate of the (meth)acrylate di-ammonium salts according to the invention was quite high and that practically no residual cationic monomers were detected after 3 hours of polymerisation. The acrylates were more reactive compared to the methacrylates. The polymerisations proceeded smoothly with good temperature control.

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The obtained polymers have a high molecular weight, generally at least 7.000.000g/mol (based on intrinsic viscosity measurements in 0,5 mol/l NaCl at 25°C).

The emulsions obtained had a high percentage offactive material (40 wt%) and were free of coagulum.

It was found that more than 90 % of the cationic monomers were accessible in the polymer.

The polymers according to the invention were therefore very useful for different purposes, and in general for whatever aqueous solid-liquid separations are required. They can be used in ion-exchange resins, for coatings, in personal care products, in cosmetics, as fabric softener, as bloode, as coagulant, as dye fixative, in oil field chemicals, in dispersants or as surfactants. They can also be used for fine retention in paper making, as flocculants, in the purification of municipal and industrial waste waters, in mines, quarties and drilling muds, in the assisted recovery of petroleum and in drinking water treatment, as stabilisers for emulsion polymerisation and in pharmaceuticals.

The polymers according to the invention showed good results when used as flocculants, especially for the treatment of industrial and municipal waste water. The present invention also relates to the use of a polymer according to the invention as flocculant.

The invention is further illustrated by tile following examples:

The following abbreviations are used:

AM: acrylic acid methylester

MAM: methacrylic acid methylester

25 MeOH: methanol

MeCl: methylchloride

BzCl: benzylchloride

BDMAP: 1,3-bis(dimethylamino)-2-propagiol

BDMAPA: 1.3-bis(dimethylamino)-2-propyl acrylate

BDMAPMA: 1.3-bis(dimethylamino)-2-propyl methacrylate

BDMAPA.2MeCl: 1.3-bis(trimethylaminonium)-2-propyl acrylate chloride

BDMAPMA.2MeCl: 1.3-bis(trimethylaminonium)-2-propyl methacrylate chloride

BDMAPA.MeCl: 1-(dimethylamino)-3-(trimethylaminonium)- 2-propyl acrylate chloride

BDMAPMA.MeCl: 1-(dimethylamino)-3-(trimethylaminonium)- 2-propyl methacrylate

35 chloride

BDMAPA.2BzCl: 1.3-bis(benzyldimethylammonium)-2-propyl acrylate chloride
BDMAPMA.2BzCl: 1.3-bis(benzyldimethylammonium)-2-propyl methacrylate chloride
BDMAPA.BzCl: 1-(dimethylamino)-3-(benzyldimethylammonium)- 2-propyl acrylate
chloride

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BDMAPMA.BzCl: 1-(dimethylamino)-3-(benzyldimethylammonium)- 2-propyl acrylate chloride

MEHQ: methyl ether of hydroquinone

PTZ: phenothiazine

5 TNPP: tris(nonylphenyl)phosfite

DPPD: N,N-diphenyl-1.4-phenylenediamine

EDTA: ethylenediaminetetraacetic acid

DBTO: dibutyltinoxide

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Example 1 : Synthesis and purification of 1,3-Bis(dimethylamino)-2-propyl methacrylate

The transesterification reaction is performed in a 3 litre jacketed glass reactor (Sovirel), equipped with a stirrer, temperature prope, air sparge and an adiabatic column filled with 7 structured metal gauze packings (Sulzer type DX). Distillation rate is controlled by a temperature controlled reflex set-up, with water-cooled 15 condenser, on top of this column. The reactor is foaded with 1950 g MAM (19.5 moles). 1.4 g MEHQ, 1.4 g PIZ and 1.4 g TNPP. This mixture is heated to reflux (96 °C), at atmospheric pressure and 95 g MAM confaining traces of water are distilled off. 4.5 g dry Li₂O (0.15 moles) is added. The reactories heated with oil, at a temperature 25 °C higher, compared to the reaction mixture, and 3 litres/hour air is sparged 20 through a sintered metal diffuser. Over a period of 100 minutes 447 g BDMAP (3.06 moles) are introduced. The methanol produced by the transesterification is distilled of at 78-94°C at a reflux ratio 5/1-15/1. In total 755 g MeOH/MAM is collected containing 97 g (3.03 moles) MeOH. The reaction takes about 6 hours. Mass temperature in the reactor reaches 108-110°C affithe end. Part of the excess MAM is 25 distilled to eliminate the last traces of MeOH. The cooled reaction mixture is passed on a Buchner filter covered with a thin layer of silica (Merck Silicagel 60) to remove the catalyst. At this stage the conversion of BDMAP \$ 98% and the yield of BDMAPMA 95 %. (determined by GC). MAM is removed by distillation at a maximum boiler temperature of 65 °C, by increasing vacuum from 20 up to 2 kPa. The obtained 30 reaction mixture (663 g) is purified in a fractional distillation set-up, equipped with a column containing 16 structured SS packings (Sulzer DX) at 0.5 kPa. A top fraction of 31 g is obtained between 60-91.5% (reflux ratio 10/1) containing BDMAP and BDMAPMA to be recycled. Heart cut 518 g (2.4 moles) (> 99.5% pure) BDMAPMA is distilled at 91.5°C (reflux ratio 1/1); the bottom emperature increases from 105°C till 35 110°C.

The collected product is stabilised with 500ppm MEHQ.

The distillation residue (102 g) still contains about 60 % BDMAPMA.

Examples 2 to 6 and comparative examples 7 to 14:

The transesterification, described in example 1, was repeated in other conditions and with a series of other catalysts. The results are shown in Table 1.

Table 1:

IBDIO				<u>gi</u>	<u> </u>		И			
No	Catalyst	Weight	MA	V1 /		Max	Ш	Conv.	Molar	Sum of
		% on	BD	MAP		T	Ш	BDMAP	yield	Michael
}]		BDMAP	(mo	lar)		(°C)		(%)	BDMAPMA	Addition
1.	•	•	Add	ition					(%)	Products
			mo	de*						% molar
1	Li ₂ O	1	6	E .		110		98	95	2
2	Li ₂ O	Q.5	5	В		107	111	85	81	4
3	Li ₂ O	2	5	<u>ç</u> :		112		98	85	8
4	Li ₂ O	1 .	1	В		110	1	95	90	5
5	LiOCH3	1	5	B.		109	1	94	80	10
6	LiOH .	1	6	₱;	П	110		95	85	6
7	NaOCH3	1	6	B		109	1	11	8	nd
8	Ca-acetyl-	1	5	В,		107	1	0	0	nd -
	acetonate			, .	П				·	
9	DBTO	1.5	5	B		10	1	0	0	nd
	Tegokat23			4				u P		
10	DBIO	1.5	5	B	П	10	ð	5	2	nd
	Tegokat 248	1 .		3 :	П	1		1		
11	Octabutyl-	1	5	В		10	1	0	0	nd
1	tetrachloro				П		H			ı I
	_d1-	ļ];	П					
	stannoxane	·	\cdot] !						
12	Dibutyl-	1	5	B		10	7	0	0	nd
	dioctyl			9					j	l l
	tetrachloro-]						1
	di-			1						<u> </u>
1	stannoxane			4					1	
13	Titanium-	1	5	В	1	1:	9	0	0	nd
	(IV)ethoxide			4:						
14	Titanium-	1	5	В	1	1.	Ö	0	0	nd
	(IV)-] '						· .
	isopropoxide			4		G				
					_					

5 (*Mode C = Continuous addition of BDMAP; Mode B = BDMAP is loaded batch-wize.

nd : not determined)

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80 °C.

Only the Lithium containing catalysts proof to be good transesterification catalysts.

Almost no or no reaction was observed when using sodium methylate. alkyltitanates or tin catalysts at temperatures below 115 °C. At higher temperatures polymerisation occurred despite the addition of inhibitors.

Example 15 : Synthesis and purification of 1,3-Bis(dimethylamino)-2-propyl acrylate

In the same equipment as example 1, 2000 g (20 moles) Ethylacrylate. 1.4 g MEHQ, 1.4 g PTZ and 1.4 g TNPP are introduced. 3 l/h air sparge is installed. Traces of water are removed by distillation of 120 g ethylacrylate at 80°C/75 kPa. 4.5 g dry Li₂O (0.15 moles) is added. Over a period of 2 hours, 447 g BDMAP (3.06 moles) are introduced. The ethanol produced in 3 moles) is distilled off together with part of the ethylacrylate at a temperature, between 66-80 °C/75 kPa at top of the distillation column, maximum temperature in the boiler reaches 102°C. After 11 hours the mixture is cooled down and filtered over silica. At this stage the conversion of BDMAP is 93 % and the yield of BDMAPA reaches 87 % (determined by GC). 1.5 g DPPD is added, and ethylacrylate is distilled off at reduced pressure (20 to 2 kPa). 750 g reaction mixture containing 530 g BDMAPA is distilled in the same fractional distillation set-up as described in example 1, at 0.4 kPa.

426.5 g (2.13 moles) >99 % pure BDMAPA (GC) is obtained as heart cut at 0.4 kPa and

Examples 16 to 17 and comparative examples 18 to 21:

The transesterification, described in example 15, was repeated in other conditions and with a series of other catalysts. The results are shown in Table 2.

No Catalyst. % Alkyl- Hours Max Conv. Molar Mich. weight on acrylate Mode T (°C)/ BDMAP yield Addit BDMAP BEBMAP in kPa % % %	
BDMAP in kPa % % %	ıael
BEMAP in kPa % % %	ition
	lucts
15 Li ₂ O 1 % Ethyl 11 C 102 75 93 87 6	
16 LIOCH ₃ Ethyl 11 C 100/75 97 77 12	
1.25%	
17 LiOCH ₃ 2% Methyl 5 B 80/80 95 60 30	
18 BaO 2% Ethyl 4 B 97/75 85 18 25	
19 NaOCH3 1% Methyl 6 B 90/101 0 0 nd	

				ı	1	111			
ĺ	20	Ca-	Methyl	6	В	87/101	0	0	nd
		acetylacetonate			1				
		2 %							
.	21	DBTO Tegokat	Ethyl	6	B	98/77	0	0	nd
		23 2 %		1				ļ	

(*Mode C = Continuous addition of BDMAP; Mode B = BDMAP is loaded batch-wize)

nd : not determined

Examples 22-23: Synthesis of 1.3-Bis(dimethylamino)-2-propyl acrylate under pressure

To obtain reaction temperatures above 100 °C, the transesterification of AM with BDMAP, such as described in example 15 with ethylacrylate, was repeated under pressure. The results are shown in Table 3. The equipment described in example 1 was constructed in 316 SS for these trials. BDMAP was added by means of a dosing pump.

Table 3:

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No	Catalyst %	Acryl	Hours	Max	Conv.	Molar	Michael
	weight on	ate	Mode	T(°C)/	BDMAP	yield	Addition
	BDMAP		feed*	Press in		BDMAPM	Products
			BDMAP	kPa		A	
22	Li ₂ O	AM	5 C	103/135	92	53	25
	1%			1 .			
23	LIOCH3	AM	6 C	104/135	95	61	20
\	1.5 %	_			8		

[*Mode C = Continuous addition of BDMAP; Mode B = BDMAP is loaded batch-wize]

Examples 24 to 28: Reaction of BDMAPMA with methylchloride

The quaternization reactions 24 to 28 are performed in a 2 litre, jacketed glass autoclave (Büchi AG), equipped with a stirrer, plunger, temperature and pressure probe and equipped with a bottom valve.

In example 28, to 193 g (0.9 moles) BDMAPMA in 800 g acetonitrile, 115 g (2.27 moles) methylchloride are added over a period of one hour. The mixture is heated up

A white solid starts to precipitate. After 23 hours at 80-82°C the mixture is cooled to room temperature, the excess of MeCl is stripped with nitrogen and the solid is filtered over a pressure filter. After drying with air and a hours under reduced pressure (10 kPa), 285 g BDMAPMA.2MeCl are obtained. (see table 4).

25 From this BDMAPMA.2MeCl, aqueous solutions can be made up to 50% weight.

These solutions contain less than 100 ppm acetonitrile (GC headspace analysis).

to 82.°C, the pressure rises to 2 bar.

Examples 24 to 27 are done according to the same procedure, except that the reaction conditions specified in Table 4 are used.

Table 4:

itrile	BDMAPMA moles	Me		Sluriy		Reaction ime	Powder		BDMAPMA .2MeCl
itrile	moles	Mo			:	ime		Yield	
	moles	Mo							.2MeCl
	moles	Mo	• [' '					
	Woles		166	ادها	P¢	(hours)	weight	%	%
			- 4	solids	1 1		(g)		
1			1	<u>' </u>	1 1	<u></u>		53.8	93.2
	0.69	1.4	15	21	50	22	136		
	0.75	7.6	14	22	60	25	183	68.7	93.1
•	0.75	1 3		<u> </u>		III	000	84.9	94.8
-	0.69	11.5	51	21 !	180	22	202	0-2.5	
702	0.03	2.0	04	126 :	85	28	277	93.1	99.4
193	0.55	_l	1 1	<u> </u>		<u> </u>	nee	972	99.0
800	0.91	2.	27	26	180	23	205		
_	793	0.75 0.69 793 0.93	0.75 1.6 0.69 1.93 2.	0.75 1.64; 0.69 1.51; 793 0.93 2.04;	0.75 1.64 22 1 21 1 21 1 21 1 21 1	0.75 1.64 22 60 0.69 1.51 21 80 793 0.93 2.04 26 85	0.75 1.64 22 60 25 60 60 60 60 60 60 60 6	0.75 1.64 22 60 25 183 0.69 1.51 21 80 22 202 793 0.93 2.04 26 85 28 277	0.75 1.64 22 60 25 183 68.7 0.69 1.51 21 80 22 202 84.9 793 0.93 2.04 26 85 28 277 93.1 0.69 0.97 0.65 1.80 23 285 97.2

Example 29-31 : Reaction of BDMAPMA with methylchloride and recycling the filtrate

The quaternization reactions 29 to 31 are performed in a 10 litre, SS Hofer autoclave in the same reaction conditions as example 28. The filtrate is reused in the following reaction.

Table 5:

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Table	e 5 :			- 1	11		ι',	-1	•			
		Aceto	BDMAPMA	Me	·Cl	Slura	γÌΤ	ì	React	Powder	Molar	Purity
	Recycle	nitrile	POWEY 14112		1	1		41	time		Yield	BDMAPMA. 2MeCl
	number	σ	moles	M	oles	% .			(hours)	weight	%	%
Dr.	TI CHILD	5				solid	s.	9		(g)		
100		3923	3,50	8.	75	211	7	2	22	1037	90.2	98.0
29	<u> </u>		3.50	B	75	22	-	71	23	1095	96.0	99.5
30	Rec 1	3884			<u>' 1</u>		14		24	1113	98.6	99.6
31	Rec 2	4015	3:50	IR	75	22		_	-	1110		
<u> </u>					1 .1	!;			lii :			

Examples 32 to 34: Quaternization of BDMAP(M)A with benzylchloride or methylchloride 15

Comparable results were obtained for the benzylchloride / BDMAPMA: methylchloride/BDMAPA and benzylchloride/BDMAPA reactions in the 21 reactor following the procedure of example 28.

The results are shown in Table 6.

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Table 6: Quaternization of BDMAP(M)A with benzylchloride or methylchloride

				P 11		सा				
Γ	Aceto	R-Cl		(meth)actyl-	Slutry	1011	React	Powder	Molar	Purity
	nitrile			ate			time		Yield	
ш	g	1	Moles	moles	%	₽ď		weight	96	%
l	·		•	(.)	solids	G	hours	(g)		
32	800	MeCl	1.72	BDMAPA ·	18 [80		205.0	97,7	BDMAPA.
]		ŀ		0.69						2MeCl
					1					99.5
33	1000	BzCl	1.45	BDMAPMA	20:	80	24	250.0	96,5	BDMAPMA
.	1	ł	1	0.56				}		.2BzCl
	<u> </u>					. 1				99.6
34	1000	BzC1	1.45	BDMAPA :	20	80		245.0	96	BDMAPA.
1	{]		0.56	2					2BzCl
				3.0	(1			99.1

Comparative Examples 35 and 36

For trials 35 and 36 the quaternization was realised in chloroform following the procedure described in patent WO 01/55089. The formed salts were extracted with water to form a -50 % aqueous solution.

Table 7: Quaternization of BDMAPMA with benzylchloride or methylchloride in chloroform

		-			1 4 1 4			#1	, , , , , , , , , , , , , , , , , , , 		
1	Chloro	R-Cl		BD-	Aq.	T	React	Press.	Molar	Purity	
	lorm			MAPMA	sol.		time		Yield	9/	6
m	g _.		Moles	moles	%	C	(b)	kPa	%		
35	1234	MeCl	3.50	1.75	50	50,	25	300	72.0	BDMAPMA	BDMAPMA
						1 :			Ì	.2MeCl	.MeCl
] (75.4	24.6
36	1414	BzCI	4.0	2.0	48	52	25	100	84.7	BDMAPMA	BDMAPMA
ł	1					1				.2BzCl	.BzCl
					1.1					89.5	10.5

In our attempts to perform the quaternization with MeCl or BzCl in aqueous solution, partial hydrolysis was observed due to the longer fraction time (12-24 hours) for the quaternization of the second amine function. Pure bisquaternized products in solution could not be obtained and it, is almost impossible to purify the obtained mixtures.

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Table 8 : Comparison of the impurities/purity of 50 wt% bisquaternized methacrylate monomer solutions in water

Q	OUS III Marci			W1	
ſ	Bisquaternized	R-C1		i.	methacrylic
١	product from			% weight	acid
	trial		١.	P1	% weight
ļ	31	MeCl		-	0.1
	33	B¢C1		< 01	< 0.05
	35	MeCl		-	2.5
	36	BzC1		0.3	1.3

Example 37: Inverse emulsion copolymerisation of BDMAPMA.2MeCl and acrylamide

This procedure is typical for the synthesis of 1 kg copolymer emulsion with 40 % active material, containing 25 % BDMARMA.2NeCl

Preparation of the aqueous phase; to 300 g acrylamide dissolved in 182 g

demineralized water. 200 g of a 50% aqueous solution of BDMAPMA.2MeCl. 0.2 g EDTA and 0.2 g potassium bromate were added. After stirring for 30 minutes 0.7 g

lactic acid and 5 g adipic acid were added (pH=3.5).

Preparation of the oil phase: in a 2 litre S5 beaker equipped with mechanical stirrer

240 g Isopar M (Isoparaffinic solvent, ExxonMobil), 18 g Montane 70 (Sorbitan mono isostearate, Seppic) and 4.6 g Atlas C-1086 (Polyoxyethylene sorbitol hexaoleate.

Uniqema) were mixed.

The aqueous phase was transferred duickly to the oil phase under stirring. The mixture was emulsified for 30 seconds at 8000 rpm using a homogenizer.

Polymerisation: the resulting emulsion was introduced in a 1 litre explosion proof CEMCO reactor equipped with a 3 blade axial flow impeller.

The emulsion was continuously purged with a nitrogen flow (1.5 l/min) for 45 minutes. At 40 °C a 0.5 g solution of 0.2 glazobis 2,4-dimethylvaleronitrile) catalyst in xylene was added through a septum on top of the reactor. The nitrogen flow was limited to 1 ml/min. The reaction was maintained at 40°C for 2 hours, and then increased up to 48°C in a time span of 3 hours, with reintroduction of 0.12 g catalyst after 3 hours and 4 hours of reaction After 5 hours, 1.2 g of sodium metabisulfite in 3 g water was added and the reaction temperature was increased to 55 °C for 1 hours.

The resulting emulsion was free of acrylamide (less then 100 ppm), showed no coagulum content and had an intrinsic viscosity (IV) of 11dl/1. (See table 9)

Examples 38-46

The same procedure of example 37 was followed with varying amounts of acrylamide and monomers BDMAFMA 2MeCl or BDMAPA.2MeCl to obtain emulsions containing 40% active material. The results are represented in Table 9.

Terpolymers were also prepared in Examples 45 and 46 in the same way by using MAPTAC (Röhm) (methacrylamidopropyltripethylammoniumchloride) and AOETAC (2-(acryloyloxy)ethyltrimethylammoniumchloride, Mitsui Chemicals) as comonomers.

10 Table 9

5

No	Cationic	Initial	П	Inverse	Pr	duct	Cationic
1	monomer	conc %	$ \cdot $	viscosit	vis	cosity	monomer in
1				d1/4	сP		polymer
			.				%
37	BDMAPMA.2MeCl	25	П	11.0	52	9 kg	23.2
38	BDMAPMA.2MeCl	5	1	11.8	56	0	4.8
39	BDMAPMA.2MeCl	10		10.4	56	0	9.4
40	BDMAPMA.2MeCl	40		11.1	38	0	35.7
41	BDMAPA.2MeCl	5		12.6:	40	0	4.8
42	BDMAPA.2MeCl	10		14.1	68	0	9.4
43	BDMAPA.2MeCl	25		10.3	54	0	23.1
44	BDMAPA.2MeCl	50	1:1	16.8	54	0	45.0
45	BDMAPA.2MeCl	5	Π	11.4	56	0	nd .
	MAPTAC	5					1
46	BDMAPA.2MeCl	5	П	10.7	56	0	nd
	AOETAC	20					

nd = not determined

The polymer obtained in Example 37 was tested for flocculation properties in a standard test with a kaolin suspension (6:6 g/l) and compared to commercial cationic polymers Alpinefloc™ E1+(−50 % canonic monomer).

The turbidity of the filtrate was measured as a function of the polymer concentration.

The results obtained at pH 7.02 and pH 8, respectively, are represented in Figures 1 and 2. In these figures, the results obtained with the polymer of Example 37 are labelled •: the results obtained with Alpinefoct E1+ are labelled .

These results show that better performances are obtained with the polymer of

20 Example 37 with respect to the polymer concentration and the transparency of the filtrate than with commercial polymers containing more cationic monomers.

Claims

5

(meth)acgulate di-ammonium salt of formula Process for the manufacture of 1. W.

$$H_{2}C = C$$

$$CH_{2} = N^{+} R^{2}$$

$$R^{2}$$

$$R^{2}$$

$$R^{2}$$

$$R^{2}$$

$$R^{2}$$

$$R^{2}$$

$$R^{3}$$

$$R^{2}$$

$$R^{2}$$

$$R^{3}$$

$$R^{2}$$

$$R^{2}$$

$$R^{3}$$

wherein \mathbb{R}^1 represents hydrogen or methyl, each \mathbb{R}^2 , independently, represents an alkyl comprising from 1 to 4 carbon atoms, each R3, independently, X , independently, represents an represents an alkyl or an aralkyl and each anion, comprising

(meth)acrylate of formula (II) (1) the reaction of the di-aming

$$H_{2}C = C$$

$$G = CH_{2}$$

$$H_{2}C = C$$

$$G = CH_{2}$$

$$H_{2}$$

$$G = CH_{2}$$

$$H_{2}$$

$$G = CH_{2}$$

$$H_{2}$$

$$G = CH_{2}$$

$$H_{2}$$

$$H_{2}$$

$$H_{2}$$

$$H_{2}$$

$$H_{3}$$

$$H_{4}$$

$$H_{2}$$

$$H_{3}$$

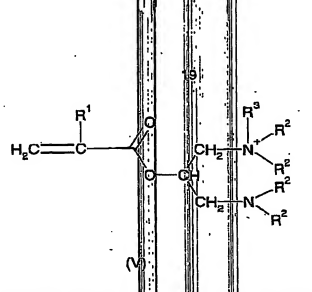
$$H_{4}$$

$$H$$

with at least 2 equivalents of at least one likyl or aralkyl derivative of formula R³X in an organic solvent containing at most 5000 ppm of water and wherein the compound of formula (I) has a solubility at 25 °C of less than 1 g/100 g of 15 solvent and wherein the solubility of the corresponding amino-(meth)acrylate animonium salt of formula

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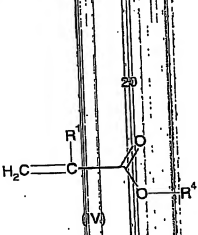
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has a solubility at 25 °C of at least 20 g/100 g of solvent; and
(2) the separation of compound of formula (1) from the reaction mixture without dissolving it in water.

- 2. Process according to claim it wherein the solvent is accionitrile.
 - 3. Process according to claim 1 or 2, wherein the reaction is effectuated at a temperature of 40 to 100 °C :
 - 4. Process according to any of claims 1 to 3, wherein the separation of the (meth) acrylate di-ammonium salt of formula (1) from the reaction mixture is done by filtration.
- 15 5. Process according any of claims 1 to 4, wherein the reaction mixture obtained after step (2) is recycled.
- 6. Process according to any of claims 1 to 5, wherein the di-amino (meth)acrylate of formula (II) used in step (I) is prepared by the transesterification of a 1.3-di-amino-2- propanol of formula (III)

wherein each \mathbb{R}^2 , idependently represents an alkyl comprising from 1 to 4 carbon atoms, with a (meth) acrylate of formula (IV)



wherein R⁴ represents an alkyl comprising from 1 to 4 carbon atoms, in the presence of a lithium-catalyst.

- 5 7. Process according claim 6. wherein the lithium catalyst is chosen from Li₂O, LiOCH₃, LiOH and their mixtures.
 - 8. Process according to claim 6 or 7, wherein the transesterification is done at a temperature not exceeding 120 °C.
 - 9. Process according to any of the preceding laims, wherein R¹ is methyl.
 - 10. Process according to any of the preceding claims, wherein R² is methyl.
- 15 11. Process according to any of the preceding claims, wherein R³ is methyl.
 - 12. Process according to any of claims it to 10 wherein R³ is benzyl.
 - 13. Process according to any of claims 6 to 12 wherein R4 is methyl.
 - 14. Process according to any of the preceding claims, wherein X is Cl.
 - 15. (Meth)acrylate di-ammonium salt of formula (I) obtainable according to the process of any of claims 1 to 11, being in solid form.
 - 16. (Meth)acrylate di-ammonium salt of formula (I) obtainable according to the process of any of claims 1 to 14. comprising less than 10 mole % relative to the amount of (meth)acrylate di-ammonium salt of formula (I) of the corresponding amino-(meth)acrylate ammonium salt of formula (V).
 - 17. Use of the (meth)acrylate disammonium salt according to claim 15 or 16 for the manufacture of a polymer comprising less than 10 mole % of units derived

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from an amino (meth)acrylate ammonium salt of formula (V).

- 18. Polymer obtained from a monomer composition comprising, per 100 parts by moles.
- 5 (a) from 1 to 100 parts by moles of at least one (meth)acrylate di-ammonium salt of formula (i) according to claim 15 or 16.
 - (b) from 0 to 99 parts by moles of a least one acrylamide monomer of formula

 [VI)

wherein R⁵ is hydrogen or methyl. R⁶ and R⁷ are, independently, hydrogen, alkyl comprising from 1 to 6 carbon atoms, optionally substituted by one or more hydroxy or alkoxy groups;

and less than 10 mole % relative to the amount of (meth)acrylate di-ammonium salt of formula (i). of amino imeth)acrylate ammonium salt of formula (V).

19. Use of a polymer according to claim 18 as flocculant.

ឲ្យប33

dimethlacrylate di-ammonium salts The invention relates to the responding to formula (1)

wherein R1 represents hydrogen drimethyl earli R2, independently, represents an alkyl comprising from 1 to 4 carbon atoms: each 189, independently, represents an alkyl or an aralkyl and each X , independently, represents an anion; having a high purity and their use as monomers for the synthesis of polymers useful

as cationic flocculants in water tre 10

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